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# Formation of cyanide compounds during preparation of gold surfaces evidenced by surface-enhanced Raman spectroscopy

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## Abstract

We report surface-enhanced Raman spectroscopy (SERS) measurements on nanostructured gold surfaces prepared using a primary vacuum sputtering technique. Even if no molecules have been deposited on the metallic surface, a clear Raman signal is observed. A very intense and sharp peak around  $2,100\text{ cm}^{-1}$  together with low-wavenumber modes are visible. The comparison between the experimental spectra and those obtained using density functional theory calculations confirms that these new modes are associated with the cyanide compound  $\text{Au}(\text{CN})_2^-$ . This proves the high sensitivity of SERS for the chemical analysis of metallic surfaces. It also shows that the pollution should be carefully taken into account in SERS analysis. We also demonstrate that high power laser illumination transforms these toxic compounds into carbon species; this method can be used to clean SERS substrates.

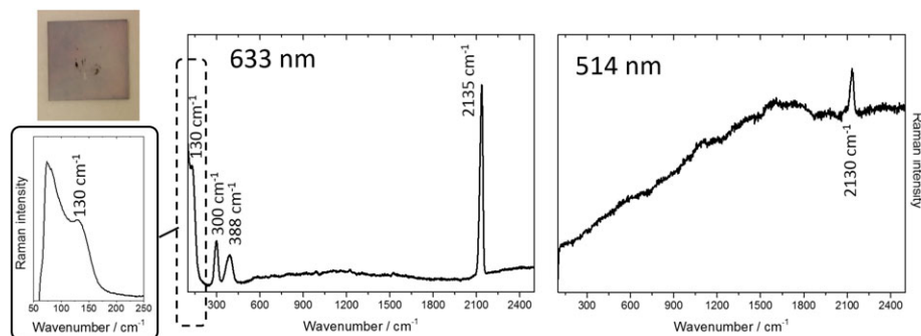
## KEYWORDS

cyanide compounds, gold, surface-enhanced Raman spectroscopy

## 1 | INTRODUCTION

Surface-enhanced Raman spectroscopy (SERS) is an analytical method based on classical Raman spectroscopy associated with optical nano-antenna.<sup>[1]</sup> It combines the advantage of Raman spectroscopy, a vibrational spectroscopy with a very low cross section, with the high sensitivity of enhanced optical processes. For most SERS studies, optical nano-antenna is based on metallic nanostructures with strong plasmonics resonances. It is well established that if the wavelength of the optical excitation corresponds to a plasmonic resonance of the metallic nanostructures, a strong enhanced electromagnetic (EM) field is produced in the near field of the antenna. The places where the EM field is enhanced are called hot spots. If a molecule is localized in such a hot spot, which usually

occurs for those with a strong chemical affinity for metallic surfaces, its Raman cross section can be increased by several orders of magnitude. This is the basic principle of SERS. The process is so efficient that single-molecule detection with SERS has been reported.<sup>[2–4]</sup> In addition, SERS offers several advantages: It is a fast, (acquisition time takes typically a few seconds, a couple of minutes if the signal intensity is low) easy, and nondestructive analysis tool. As a consequence, SERS is now currently used in a large field of applications, notably for the detection of pollutant.<sup>[5,6]</sup> For instance, for the analysis of ambient air composition, Mosier-Boss and Lieberman<sup>[7]</sup> have used an SERS sensor based on silver chemically functionalized substrates mounted on a thermoelectric cooler to detect volatile organic compounds. With their method, they successfully detected various compounds such as chlorinated



**FIGURE 1** Typical Raman spectra of blank surface-enhanced Raman spectroscopy surface (633- and 514-nm excitation) [Colour figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

solvents, aromatic compounds, and methyl *t*-butylether. Thus, SERS appears as a privileged tool to study air pollution.

Usually, the protocol to perform SERS is divided in 3 steps: (1) preparation of the metallic (usually gold or silver but any metals with plasmonic resonances in the visible can be used) nanostructured substrate, (2) deposition of the molecule on the SERS substrate (most of the time, this process is done by immersion in a low concentration solution followed by a thorough rinsing to remove molecules in excess, and (3) Raman measurement is performed using a classical Raman spectrometer. Surprisingly, even if SERS is regarded as an extremely efficient and sensitive process, most of SERS studies are performed under ambient air conditions. This experimental procedure has two major drawbacks. First, the metal can be oxidized losing its optical properties. This is particularly true for silver: It is well known that silver SERS substrates must be used immediately after preparation otherwise the chemical degradation of its surface will cause a rapid decrease of the SERS signal.<sup>[8]</sup> This problem can be avoided using gold, which is more chemically stable compared with silver. Second, in ambient conditions, the substrate can be easily polluted by molecules present in the air. Nevertheless, in most cases, molecules studied by SERS have a strong chemical affinity for metallic surfaces and their adsorption removes the possible contaminants. At the end, the SERS signal corresponds to the Raman signature of the molecule, except for specific modes associated with the interaction between the molecule and the metal.

Nevertheless, the question of the SERS signal of the nanostructured metallic surface when no molecules is deposited remains open. In such conditions, polluting molecules should still be present, and indeed, it can be found in scientific forums on the internet discussions about Raman modes observed on blank gold surfaces.<sup>1</sup> The origin of such

spectral signatures remains unclear and may overlap with compounds of interest while analysis.

In this study, we have performed SERS measurements on a blank gold SERS substrate. Surprisingly, we have observed Raman signature of cyanide compounds. It proves the adsorption on the gold surface of chemical species.

## 2 | METHODS

Gold nanostructured surfaces were prepared using metallic sputtering under primary vacuum ( $10^{-1}$  Pa). We used a Jeol Fine Coat ion sputter JFC-1100 with a voltage of 1.3 kV. The deposition rate was set at 10 nm/min. Before sputtering, the glass slides were washed with alcohol and we have systematically used gloves to avoid any contamination. A typical picture of a glass side after gold deposition can be seen in Figure 1. Its typical color is between blue and purple.

Raman experiments were performed using a Horiba Jobin Yvon HR800 Raman spectrometer associated with a 600-lines/mm array. This device gives a spectral resolution better than  $0.5 \text{ cm}^{-1}$ . SERS measurements were performed with a laser power of 0.1 mW for the 633-nm excitation and 0.35 mW for the 514 nm. The laser spot was focused on the sample surface using a 100× objective (NA: 0.9). The scattered light was collected through the same objective in a backscattering geometry. Time acquisition was optimized to ensure a high signal to noise ratio and was typically a few minutes.

DFT calculations were performed using Gaussian09 suit of programs.<sup>[9]</sup> They were carried out in the DFT formalism. We used the B3LYP functional and D95V basis set for C and N, whereas LANL2DZ one was used for Au atoms. Au, C, and N atoms of  $\text{Au}(\text{CN})_2$  molecule were kept free to move whereas those of Au layer were blocked in a monolayer structure, in order, in a first rough approximation, to mimic Au surface. The aim of this protocol was to check  $\text{Au}(\text{CN})_2$  bending ability.

<sup>1</sup>[https://www.researchgate.net/post/What\\_is\\_the\\_peak\\_at\\_around\\_2100\\_cm-1\\_on\\_a\\_clean\\_gold\\_surface\\_due\\_to](https://www.researchgate.net/post/What_is_the_peak_at_around_2100_cm-1_on_a_clean_gold_surface_due_to)

### 3 | RESULTS AND DISCUSSION

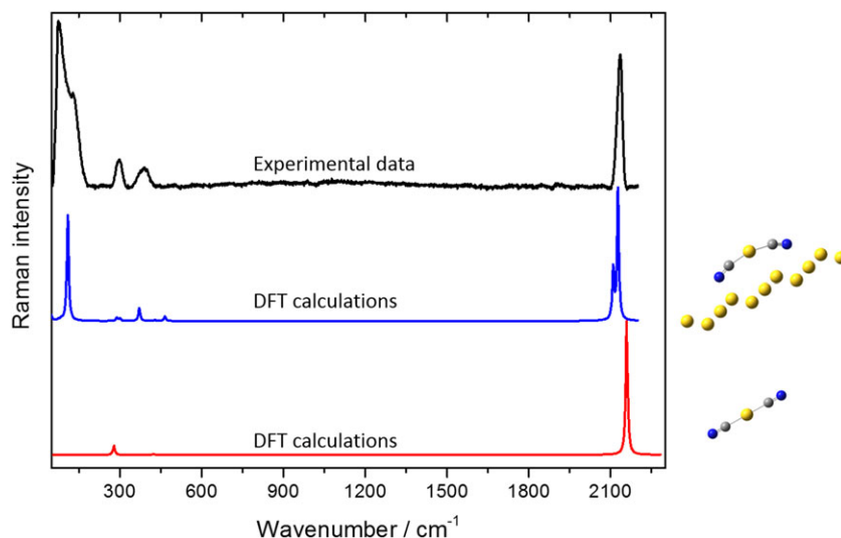
Figure 1 depicts a typical Raman spectrum of a blank SERS sample. As no molecules have been normally deposited on the gold surface, we should not observe any Raman modes. Surprisingly, this is obviously not the case: With the 633-nm excitation, four modes are visible. A very intense mode is observed at  $2,135\text{ cm}^{-1}$ , two low-wavenumber modes with a similar intensity are observed at  $300$  and  $388\text{ cm}^{-1}$ , and a last mode is visible at  $130\text{ cm}^{-1}$  in the shoulder of the Rayleigh scattering (evidenced in the inset of Figure 1). It must be said that we have observed a remarkable stability and homogeneity of this Raman signature all over the sample. We performed measurements on the same sample 4 months later, and the signal was still the same. This point is remarkable as, for SERS measurements (in particular in the case of single-molecule detection), strong fluctuations of the Raman signal are usually observed.<sup>[10]</sup> This is not the case here; the signal is remarkably stable. In addition, we performed this measurement using two Raman spectrometers in different laboratories, and the signal was still the same.

To be sure that we really have an SERS effect, we performed Raman measurements using a different excitation wavelength: 514 nm instead of 633 nm. It is established that the EM enhancement of an SERS substrate is strongly related to its plasmonic properties: It is much higher when the excitation wavelength is close to the plasmonic resonance.<sup>[11]</sup> For gold substrates prepared with our protocol, the resonance is much closer to 633 than 514 nm, which is confirmed by optical absorption measurements (not shown). As a consequence, the EM enhancement should be much higher at 633 nm compared with 514 nm. As can be seen, comparing both excitation wavelengths in Figure 1, this is exactly what we observed: At 514 nm, the Raman signal, even if it is still present, has a very low intensity. Low-wavenumber modes are not visible anymore, and only the  $2,130\text{ cm}^{-1}$  peak is still present. This result demonstrates that Raman signal is really coming from an SERS effect. It must be noticed that we also observe a rather strong background with the 514-nm excitation, this probably comes from the luminescence of gold nanostructures.<sup>[12]</sup>

The question now is the identification of the chemical species responsible for this SERS signal. Bulk gold has no Raman signal, and the unidentified modes cannot be associated with the vibrational signature of this metal. In the particular case of gold nanoparticles, acoustic modes can be observed at very low wavenumber (typically  $35\text{ cm}^{-1}$  for nanoparticles of 3.2 nm diameter, and the wavenumber downshifts if the diameter increases).<sup>[13]</sup> This suggests that these modes are coming from a molecule present at the surface of the metal.

There is a very large number of different kind of molecules in the air, each one having a typical vibrational signature. If all molecules are indeed adsorbed, we should observe a Raman spectrum with a very large number of modes. Obviously this is not the case here: only four sharp modes are observed. We can reasonably assume that these modes are the vibrational signature of a limited number of various molecules.

The most abundant gases present in the air are dioxygen ( $\text{O}_2$ ) and nitrogen ( $\text{N}_2$ ). The Raman spectrum of  $\text{O}_2$  has a typical band at  $1,550\text{ cm}^{-1}$ ,<sup>[14]</sup> not present in our experimental data.  $\text{N}_2$  has a band at  $2,330\text{ cm}^{-1}$ , neither observed.<sup>[15]</sup> Therefore, it appears clearly that these two gases are not responsible of the modes observed on the gold surface. As there is no spectral signature between  $1100$  and  $1800\text{ cm}^{-1}$ , it is clear that these modes do not come from carbon soot potentially present in the atmosphere.<sup>[16]</sup> In addition, it is clearly established that a molecule can give rise to an intense SERS signal only if it has a strong affinity for the metallic surface. As gold has a rather low chemical reactivity, the number of chemical functional groups that can strongly adsorb on its surface is limited. The most commonly used functional group that can react with a gold surface is the thiol function ( $-\text{SH}$ ). This kind of molecules is extensively used as probe molecules in SERS measurements.<sup>[17]</sup> In this case, some Raman bands at typically  $200\text{--}300\text{ cm}^{-1}$  appear and are associated with the Au-S stretching mode.<sup>[18,19]</sup> In our measurements, we do not observe such modes. In addition, it is also well known that the molecules with a thiol functional group have strong odors (they are used as additives in the domestic gas to facilitate its detection) and their presence in the air (even in a very low amount) can be easily detected by the experimentalists themselves. It was not the case during our experiments. We can therefore safely assume that the reported modes cannot be associated with thiol molecules. The other functional groups with an affinity for gold potentially present in the air are those with a nitrogen atom. These various compounds with nitrogen atoms can react with the gold surface to form specific molecules. It has already been shown that the CN functional group can react with gold and the SERS signature of cyanide compounds has already been extensively studied,<sup>[20–23]</sup> in particular the dicyanoaurate  $\text{Au}(\text{CN})_2^-$  molecules.<sup>[24–26]</sup> In Figure 2, we compare the DFT calculations of this molecule with our experimental Raman measurements. DFT calculations have been performed with two configurations: first, the free  $\text{Au}(\text{CN})_2^-$  molecule and, second, with the same molecule in interaction with a gold surface. It appears clearly that the molecule remains linear when it is free whereas it is bended when it bends when interacting with the gold surface. This point has already been reported<sup>[25]</sup>



**FIGURE 2** Comparison between experimental data and density functional theory (DFT) calculations [Colour figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

and is confirmed by our calculations. In this study, Jacobs et al. have also reported the transformation of dicyanoaurate ions to tetracyanoaurate ions  $\text{Au}(\text{CN})_4^-$ . We have here not observed the low-wavenumber modes associated with this new compound (notably the mode around  $500\text{ cm}^{-1}$ ); therefore, the transformation does not occur in our measurements.

Independent of the dicyanoaurate configuration, we observe a remarkable agreement between the calculations and the experimental spectra. The intense mode around  $2,100\text{ cm}^{-1}$  is notably present for both configurations, even if it is associated with a doublet with the molecule in interaction with a gold surface. The experimental peak at  $2,100\text{ cm}^{-1}$  is too broad to determine if it is composed of a doublet or a single mode. The main differences between the two configurations are for the low-wavenumber modes. For the free molecule, only one mode is observed, which can be associated with the experimental mode at  $300\text{ cm}^{-1}$ . For the molecule in interaction with the gold surface, two modes are observed in the low-wavenumber region: one at  $130\text{ cm}^{-1}$  and the other around  $390\text{ cm}^{-1}$ . Some much weaker modes are also present in the calculations, but their intensities are too weak to give an observable mode. Thus, with DFT calculations of  $\text{Au}(\text{CN})_2^-$  molecules free or in interaction with a gold surface, it is possible to identify all the experimental Raman modes. The fact that we observe both modes from the free and in interaction molecule suggest that we have different sites of adsorption on the gold surface. This is not surprising as the crystallinity of the gold nanostructures prepared using our protocol is rather poor and they offer a large variety of such adsorption sites.

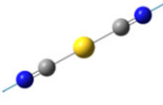
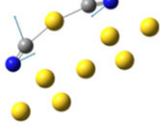
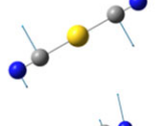
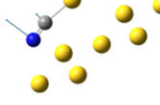
The different modes observed and calculated are reported in Table 1. The Raman signal thus originates from this cyanide molecule adsorbed on the hot spots of our SERS substrate. The high EM enhancement makes

them detectable by Raman spectroscopy even if they are present in a very low amount.

The presence of cyanide compounds on gold surfaces could be useful: A recent study performed by Kim et al.<sup>[27]</sup> have demonstrated that cyanide SERS substrates could be used to detect organic volatile compounds and metal ions. To do so, they proposed to exploit the susceptibility of the CN stretching wavenumber of cyanide on Au (around  $2100\text{ cm}^{-1}$ ). As this mode is spontaneously present in the SERS substrates (and very stable in time), our method could greatly facilitate their protocol. Nevertheless, as these molecules are highly toxic, for other applications of nanostructured gold surfaces, it is highly desirable to reduce their amount. To do so, we performed maximum laser power illumination (10 mW with the 633-nm excitation) of the sample during a few seconds. In Figure 3, we compare the SERS spectrum before and after this illumination. It is obvious that the Raman signal is notably affected. The intensity of 300, 388, and  $2135\text{ cm}^{-1}$  modes has considerably decreased, and new modes between 1100 and  $1,800\text{ cm}^{-1}$  are visible. These new modes are associated with the D and G bands of carbon species.<sup>[28]</sup> These features suggest that the cyanide molecules present at the gold surface have been partly degraded in carbon by the high power laser illumination. The catalytic activity of gold nanoparticles could also promote this reaction.<sup>[29]</sup> This method is thus efficient to reduce the amount of cyanide molecules on the surface. We have also performed SERS measurements using argon atmosphere and the Raman signature from the cyanide molecules rapidly decreased between 100 and  $200^\circ\text{C}$  confirming that they can be easily removed from the surface.

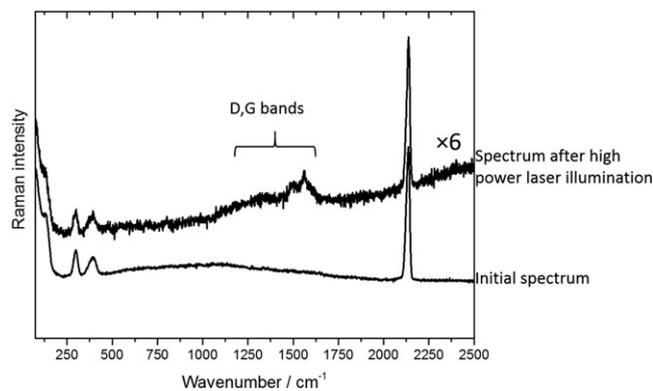
The next question is the origin of the cyanide compounds. As these molecules are not present in the air, we can reasonably assume that they are formed

**TABLE 1** List of Raman modes [Colour figure can be viewed at wileyonlinelibrary.com]

Experimental Raman frequency	DFT calculations of $\text{Au}(\text{CN})_2^-$	DFT calculations of $\text{Au}(\text{CN})_2^-$ in interaction with a gold surface	Functional mode / vibration	
2135 $\text{cm}^{-1}$ (VS)	2160 $\text{cm}^{-1}$ (VS)	2127 $\text{cm}^{-1}$ (VS) 2109 $\text{cm}^{-1}$ (S)	CN stretching	
388 $\text{cm}^{-1}$ (S)	--	371 $\text{cm}^{-1}$ (S)	Au-CN bending motion	
300 $\text{cm}^{-1}$ (S)	288 $\text{cm}^{-1}$ (S)	--	Au-CN bending motion	
130 $\text{cm}^{-1}$ (M)	--	109 $\text{cm}^{-1}$ (VS)	Au-CN bending motion	

Note. DFT = density functional theory; VS = very strong; S = strong; M = medium.

--, Not observed.

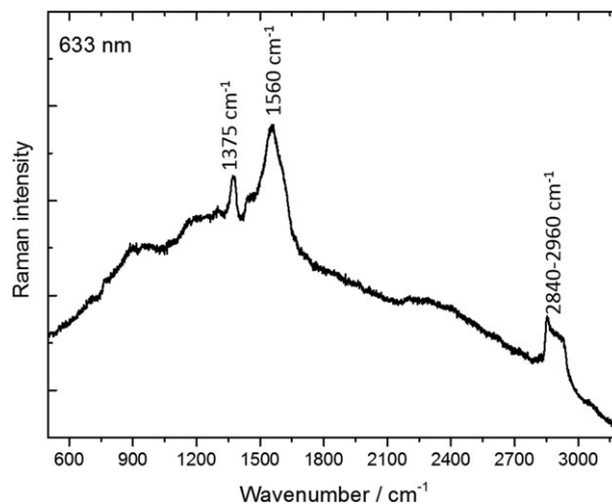


**FIGURE 3** Comparison between the Raman spectrum of the gold surface before and after high power laser illumination. Both spectra have been recorded in the same experimental conditions with the 633-nm excitation

during the sputtering process. It is clear that the use of a primary vacuum does not prevent the presence of organic molecules in the chamber that could react during the sputtering process to form the contaminant molecules on the gold surface.

We performed identical measurements using a different gold evaporator device (EDWARDS Auto 306, pressure:  $3.6 \cdot 10^{-6}$  mbar, deposition rate: 0.12 nm/s). A typical Raman spectrum of this new sample can be seen in Figure 4.

Obviously, the SERS signal is totally different. None of the previously observed modes are present, indicating that cyanide compounds are not present anymore. This result confirms that cyanide compounds are formed during the



**FIGURE 4** Surface-enhanced Raman spectroscopy spectrum of a different gold surface

sputtering process and not as a pollution coming from ambient air, otherwise the signal should be the same whatever the method used for the preparation of the gold surface.

Three modes are now observed: a first one at  $1375 \text{ cm}^{-1}$ , potentially associated with aromatic azo functions, a second one at  $1560 \text{ cm}^{-1}$ , also associated with nitrogen groups, and a broad one, between  $2840$  and  $2960 \text{ cm}^{-1}$  usually associated with C-H functions. The presence of these modes suggests that once again, nitrogen- and carbo-based molecules are present on the gold surface. In a similar way to the previous sample, if the gold surface

is submitted to a high power laser illumination, the molecules are transformed into carbon species and the D and G bands are observed (spectrum 514 here). This result proves that this method can be efficiently used to clean an SERS substrate independently of the organic species present at its surface.

In this study, we have demonstrated using SERS that organic molecules with nitrogen functions are formed during sputtering of gold under primary vacuum. For some specific conditions, these molecules can be cyanide compounds. The presence of such molecules should be systematically taken into account for the surface analysis of gold prepared using such protocols. We have also shown that an optical treatment can be used to transform these potentially toxic molecules in carbon species. This study proves the high sensitivity of SERS for the analysis of metallic surfaces.

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